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Autonomy and robustness of translocation through the nuclear pore complex: a single-molecule study

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All molecular traffic between nucleus and cytoplasm occurs via the nuclear pore complex (NPC) within the nuclear envelope. In this study we analyzed the interactions of the nuclear transport receptors kapα2, kapβ1, kapβ1ΔN44, and kapβ2, and the model transport substrate, BSA-NLS, with NPCs to determine binding sites and kinetics using single-molecule microscopy in living cells. Recombinant transport receptors and BSA-NLS were fluorescently labeled by AlexaFluor 488, and microinjected into the cytoplasm of living HeLa cells expressing POM121-GFP as a nuclear pore marker. After bleaching the dominant GFP fluorescence the interactions of the microinjected molecules could be studied using video microscopy with a time resolution of 5 ms, achieving a colocalization precision of 30 nm. These measurements allowed defining the interaction sites with the NPCs with an unprecedented precision, and the comparison of the interaction kinetics with previous in vitro measurements revealed new insights into the translocation mechanism.

Introduction

The exchange of ions, small and large molecules between cytoplasm and nucleus is mediated by the nuclear pore complex (NPC), a highly symmetrical supramolecular complex traversing the nuclear envelope (NE) (for review see Peters, 2005; Lim and Fahrenkrog, 2006; Tran and Wente, 2006; Stewart, 2007). A complex octagonal cylindrical central framework with an outer diameter of 125 nm and a length of 40 nm is covered by eight cytoplasmic filaments of ~50 nm in length and eight nuclear filaments of ~80 nm in length, forming a basket-like structure (Beck et al., 2004). The NPC is composed of ~30 different polypeptides designated nucleoporins, which yield a total mass of ~125 MDa in vertebrates (Rout et al., 2000; Cronshaw et al., 2002). A large number of nucleoporins contain repetitive sequences, the phenylalanine-glycine (FG) repeats, which are intrinsically unfolded (Denning et al., 2003).

Small molecules, which do not specifically interact with the FG repeats, permeate the NPC at rates inversely related to their Stokes radius (Keminer and Peters, 1999). The transport rates are consistent with restricted diffusion through a channel within the NPC center ~10 nm in diameter and ~45 nm in length. The translocation of molecules, which specifically interact with the FG repeats, e.g., the transport receptors importin β or karyopherin β1 (kapβ1) and transportin 1 or karyopherin β2 (kapβ2) is facilitated (Stewart, 2007). Larger substrates, containing a nuclear localization signal (NLS), do not interact directly with the NPC but bind in the cytoplasm to transport receptors of the karyopherin/importin β family. Import complexes are translocated through the NPC and dissociate in the nucleus upon binding of RanGTP. Conversely, substrates containing a nuclear export signal (NES) form ternary complexes with a transport receptor and RanGTP in the nucleus, which are translocated through the NPC, and hydrolysis of the GTP induces their dissociation within the cytoplasm. Surprisingly, passive and facilitated transport through NPCs do not interfere with each other, suggesting that the two modes of transport are most probably spatially segregated (Naim et al., 2007).

Restricted and facilitated diffusion as well as receptor-mediated transport of NLS-containing cargo molecules through...
the NPC are passive, bidirectional processes (Kopito and Elbaum, 2007). Receptor-mediated transport can proceed against concentration differences, and depends on a variety of parameters such as cytoplasmic and nucleoplasmic concentration of the cargoes, and the RanGDP and RanGTP levels in the respective cellular compartments. Probably, the mechanism by which molecules migrate through the NPC is simple thermal motion (Peters, 2007; Stewart, 2007). The FG repeats are thought to play a decisive role in the exclusion of larger molecules from the nuclear interior. Simultaneously the transport of large molecules, with which the FG repeats interact by means of transport receptors, is facilitated. However, both the spatial arrangement of binding sites within the NPC and the mechanism of exclusion of unrelated large substrate molecules are still a matter of dispute (Rout and Aitchison, 2001; Frey et al., 2006; Frey and Gorlich, 2007; Lim et al., 2007; Patel et al., 2007; Weis, 2007).

Single-molecule analysis provides unique information on spatial properties and kinetic processes that are lost by averaging over large populations of unsynchronized molecules (Zlatanova and van Holde, 2006; Ishii and Yanagida, 2007; Peters, 2007). Optical far-field single-molecule microscopy using fast and sensitive CCD camera systems is particularly suited for biological applications (Schmidt et al., 1999). Single molecules are imaged as diffraction-limited spots, and their positions can be determined with a precision in the size range of a typical protein diameter by image analysis. The localization precision that can be achieved depends predominantly on the signal-to-noise ratio (SNR), and may be as small as a few nanometers under optimal conditions (Thompson et al., 2002; Ober et al., 2004). If imaging is performed fast enough, single molecule detection allows observing the movement of single molecules even in aqueous solution (Grunwald et al., 2006a). The technique has most recently been applied to analyze the movement of single receptor and cargo molecules across the NPC predominantly in digitonin-permeabilized cells (Yang et al., 2004; Kubitscheck et al., 2005; Yang and Musser, 2006).

It might be anticipated that nuclear transport would show a quite different spatio-temporal behavior in living cells compared with digitonin-permeabilized cells. This was suggested by molecular crowding and viscosity differences in a live cell compared with digitonin-permeabilized cells, where most soluble cytoplasmic components are washed out (Banks and Fradin, 2005; Hu et al., 2007). Also, transcription is almost completely halted in digitonin-permeabilized cells, which can be deduced from the change in the intranuclear dynamics of splicing factors (Grunwald et al., 2006b), and certainly results in a significant reduction in nuclear export processes. Sustained export processes, however, should interfere with nuclear import by modifying the available intra-pore binding sites, and thus presumably, translocation times.

In this study we analyzed the translocation of single transport receptors and substrates in vivo. All experiments were based on introducing the fluorescently labeled probe molecules by microinjection into the cytoplasm of living cells. We defined the respective binding site distributions and the interaction times with the pore with an unprecedented precision. We found that all examined molecules bound in a central and spatially confined region of the NPC for a time span of roughly 5–7 ms. The interaction characteristics did not depend on the specific transport pathway. Hence, nucleocytoplasmic traffic in living cells is autonomous and robust with regard to persistent simultaneous import or export, molecular crowding, and cytoplasmic viscosity.

Results

Monitoring the association of single nuclear transport receptors and cargo molecules with GFP-labeled nuclear pore complexes

The interaction of AlexaFluor 488–labeled kapo2, kapβ1, kapβ1ΔN44, kapβ2, and BSA-NLS with the NPC was studied in living HeLa cells stably expressing GFP-POM121 (Boodor et al., 1999; Kubitscheck et al., 2005). Cells were microinjected with low concentrations of the labeled protein at room temperature. Immediately after the delivery of the probe, the equatorial plane of the nucleus was brought into focus. While illuminating a cellular region centered at the NE with 488-nm laser light at irradiance between 5.5 and 7 kW/cm², high-speed movies were acquired at a frame rate kacq of ~200 Hz. Complete and irreversible bleaching of the POM121-GFP fluorescence occurred within an illumination period of 5 s, corresponding to 1,000 single frames. The first 100 images of such an image sequence were used to determine the position of the NE. For this purpose these images were averaged yielding a single image showing the GFP-labeled NPCs with an excellent signal-to-noise ratio (SNR). After bleaching the dominating GFP fluorescence single AlexaFluor 488–labeled probe molecules became visible while moving within cytoplasm and nucleoplasm, and also upon binding to the NE. Several video stills from video sequences illustrating such events are shown in Fig. 1. In this figure, the average image illustrating the position of the NE was plotted in green (designated as: GFP image), and selected images acquired later in the movie are overlaid in red. Single probe molecules bound to the GFP-marked NE are indicated by white arrows. Fig. 1 A shows an example sequence of an experiment, where the import receptor kapo-AlexaFluor 488 was interacting with the NE, and Fig. 1 B shows an image series of a relatively long-lasting binding period of BSA-NLS-AlexaFluor 488. In separate control experiments, unrelated molecules (BSA-AlexaFluor 488 without NLS) did not show any significant interaction with the NE. Because all proteins mentioned above are involved in nucleocytoplasmic transport, we conclude that the observed binding was directly to NPCs. Thus, such image series contain information on the spatial position of the single fluorescent probe molecules with regard to the NPCs, as well as temporal information on the interaction duration of the molecules with the NPCs.

Association periods of transport receptors and transport substrates at the NPC

Individual molecules of kapo2, kapβ1, kapβ1ΔN44, kapβ2, and BSA-NLS, labeled with AlexaFluor 488 were observed for varying numbers of frames directly at the NPCs. These numbers, n, correlated with the length of the individual binding events and were translated into binding times, τ, with τ = n/kacq. These data were used to establish decay curves N(τ) quantifying the number
and Kap2, and also for an NLS substrate and Kap1 (Yang and Musser, 2006). A small deviation from a mono-exponential decay behavior was seen in the semi-logarithmic plots of Fig. 2. Although this deviation was negligible for Kap1 and BSA-NLS, a second decay component comprising a fraction of ~10% of the observed binding events was obtained for Kap2 and Kap1ΔN44 (Table I). This was in agreement with a previous study, where binding times of single transport receptors were examined in digitonin-permeabilized cells (Kubitscheck et al., 2005). Also in this system binding events were observed, which
the import event, i.e., the dissociation of kap\textsubscript{H9251} from kap\textsubscript{H9252} and release of the import substrate from the nuclear pore into the nucleoplasm (Gorlich et al., 1996).

A monoexponential fit failed to describe the decay curve of kap\textsubscript{H9252}. A two-component fit was required for a good description of the data, and yielded a short dwell time, \( \tau_{\text{kapH9252,1}} = 4.6 \pm 0.1 \text{ ms} \), and a longer time, \( \tau_{\text{kapH9252,2}} = 103 \pm 6 \text{ ms} \) (Fig. 2E, Table I).

The mutant kap\textsubscript{H9252\textsubscript{H9004N44}} has a RanGTP binding deficiency (Gorlich et al., 1996). In the association analysis this mutant showed a nearly twofold prolonged dwell time at the NPC compared with wt kap\textsubscript{H9252}. A notably extended dwell time was expected, as Ran-GTP binding is required for the termination of the import event, i.e., the dissociation of kap\textsubscript{2} from kap\textsubscript{1} and release of the import substrate from the nuclear pore into the nucleoplasm (Gorlich et al., 1996).

A monoexponential fit failed to describe the decay curve of kap\textsubscript{2}. A two-component fit was required for a good description of the data, and yielded a short dwell time, \( \tau_{\text{kapH9252,1}} = 4.6 \pm 0.1 \text{ ms} \), and a longer time, \( \tau_{\text{kapH9252,2}} = 103 \pm 6 \text{ ms} \) (Fig. 2E, Table I).

Table 1. Binding site distribution and dwell times of import receptors and model cargo

<table>
<thead>
<tr>
<th>Receptor</th>
<th>( \tau_{\text{dwell}} ) [ms]</th>
<th>Fraction [%]</th>
<th>( x_c ) [nm]</th>
<th>FWHM [nm]</th>
<th>( N_{\text{observation}} )</th>
<th>( N_{\text{trans}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>BSA-NLS</td>
<td>6.2 ± 0.3</td>
<td>100</td>
<td>-13 ± 1</td>
<td>140 ± 5</td>
<td>293</td>
<td>75</td>
</tr>
<tr>
<td>Kap\alpha</td>
<td>7.5 ± 0.8</td>
<td>100</td>
<td>-6 ± 2</td>
<td>160 ± 5</td>
<td>387</td>
<td>136</td>
</tr>
<tr>
<td>kapβ1</td>
<td>6.6 ± 0.2\textsuperscript{a}</td>
<td>100\textsuperscript{a}</td>
<td>-10 ± 2</td>
<td>160 ± 5</td>
<td>402</td>
<td>110</td>
</tr>
<tr>
<td>kapβ1\textsubscript{ΔN44}</td>
<td>11.8 ± 0.6\textsuperscript{b}</td>
<td>100\textsuperscript{b}</td>
<td>-8 ± 1</td>
<td>90 ± 2</td>
<td>244</td>
<td>60</td>
</tr>
<tr>
<td>kapβ2</td>
<td>4.6 ± 0.1</td>
<td>90 ± 1</td>
<td>5 ± 2</td>
<td>150 ± 5</td>
<td>305</td>
<td>97</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Bi-exponential fits yielded \( \tau_1 = 5.8 \pm 0.3 \text{ ms} \) and \( \tau_2 = 29 \pm 18 \text{ ms} \) with \( f_1 = 93 \pm 5 \) and \( f_2 = 7 \pm 5 \).

\textsuperscript{b}Bi-exponential fits yielded \( \tau_1 = 8.9 \pm 1.4 \text{ ms} \) and \( \tau_2 = 41 \pm 33 \text{ ms} \) with \( f_1 = 86 \pm 13 \) and \( f_2 = 14 \pm 13 \).
The requirement of two decay components for \( \text{kap}^{\beta}2 \) was in accordance with our previous study in digitonin-permeabilized cells, where we found extended interaction times of more than 100 ms for 2.5% of the observed events (Kubitscheck et al., 2005). The slow \( \text{kap}^{\beta}2 \)-dissociation was more frequent in live cells (10%) compared with digitonized cells (2.5%).

**Distances between binding sites of single probe molecules and the NE can be measured with a precision in the nanometer range**

Image sequences as those shown in Fig. 1 were used to determine both the position of the NE and the various used single fluorescent probe molecules as previously described (Kubitscheck et al., 2005). As an example, an overlay of the GFP image (background image), the corresponding deduced position of the NE (green dotted line), and the binding sites of several individual \( \text{kap}^{\beta}1\Delta N44 \)-AlexaFluor 488 molecules (red dots) deduced from one movie sequence is shown in Fig. 3 A. Approximately 30 distinct spots of GFP fluorescence were visible along the NE section of 13 \( \mu m \) length. Assuming a focal depth of 0.8 \( \mu m \), this would correspond to a pore density of \( \sim 3 \) nuclear pores per \( \mu m^2 \), which is in agreement with our previously published data (Kubitscheck et al., 1996; Kues and Kubitscheck, 2002).

During the 5-s observation time \( \sim 10 \) out of the 30 discernable NPCs were visited by \( \text{kap}^{\beta}1\Delta N44 \) molecules. These \( \text{kap}^{\beta}1\Delta N44 \) molecules were almost exclusively observed at sites of strong POM121-GFP fluorescence corresponding to labeled NPCs, whereas at NE sections devoid of GFP fluorescence almost no \( \text{kap}^{\beta}1\Delta N44 \) molecules were observed. This suggests that all NPCs are labeled by POM121-GFP. In Fig. 3 B the position of the NE (full line) and those of the observed individual \( \text{kap}^{\beta}1\Delta N44 \)-AlexaFluor 488 molecules (open circles) were plotted in a single graph. From such data the individual distances from the NPCs were determined as the shortest separation distances between the sites of receptor binding and the nearest point on the curved line indicating the NE. The distribution of these distances for \( \text{kap}^{\beta}1\Delta N44 \) molecules binding to the NE is shown in Fig. 3 C. Negative values indicate a position at the cytoplasmic face of the NPC, positive values correspond to positions on the nucleoplasmic face. The distribution has a very symmetrical shape with a mean at \( x = -3 \pm 1 \) nm. The full width at half maximum (FWHM) of the binding site distribution for \( \text{kap}^{\beta}1\Delta N44 \) of this cell is very low with only \( 80 \pm 2 \) nm.

**Transport receptors and the model transport substrate bind primarily to the cytoplasmic filaments and in the central domain of the NPC**

For all used proteins we determined the distances between observation sites and the nearest point of the NE. These distances were accumulated from all acquired measurements, and plotted in binding site histograms (Fig. 4). As above, negative values indicate a cytoplasmic position, and positive values correspond to sites on the nucleoplasmic face of the NPC. These histograms represent a novel tool to determine NPC binding sites with an unprecedented localization precision. In our previous work using

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**Figure 3.** Single \( \text{kap}^{\beta}1\Delta N44 \) molecules at the NE. For a single cell, all fitted positions (open red circles) were overlaid with the image of the NE. The position of the NE was indicated by a green line in A and B. \( \text{kap}^{\beta}1\Delta N44 \) molecules were preferentially observed at the GFP-labeled NPCs, while molecules were only very rarely observed within unlabeled regions of the NE. The fitted positions of both the NE and the single molecules were plotted to scale in B. The individual distances of the \( \text{kap}^{\beta}1\Delta N44 \) molecules from the NE were calculated as the shortest distances between the fitted position of the single molecule and the nearest point on the fit through the NPC signals marking the NE. The resulting distance distribution of \( \text{kap}^{\beta}1\Delta N44 \) molecules to the NE is shown in C. The position of the NE was defined as zero. Then, negative values described a single molecule position at the cytoplasmic face of the NPC, positive values at the nucleoplasmic face. A fit to a Gaussian yielded a maximum position of the distribution at \( -3 \pm 1 \) nm with a width (FWHM) of \( 80 \pm 2 \) nm.
two-color fluorescence, the analysis was largely limited by a lower colocalization precision caused by the two detectors and by sample instabilities (Kubitscheck et al., 2005).

Fig. 4 shows that all examined molecules bind preferably to the NPCs. The enhanced binding extends over a distance of 150 nm along the axial symmetry axis from about −80 to +70 nm with an approximate Gaussian shape. The peak positions were located at the very center of the NPC, slightly on the cytoplasmic side of the NPC (with the exception of kapβ2; see Table I). As shown previously, the tips of the cytoplasmic filaments extend ~75 nm from POM121-GFP (Kubitscheck et al., 2005), whereas the closure of the nuclear cage is presumably located at a distance of 80 nm from POM121-GFP (Beck et al., 2004). Therefore, the observed binding sites corresponded exactly to the anticipated region of the NPC, with a preference for the cytoplasmic face. Noticeably, the binding site distribution of the mutant kapβ1 ΔN44 deficient in RanGTP binding was significantly narrower than all other distributions. This may be partly due to the longer dwell time resulting in a better mean SNR of this dataset.

**Discussion**

Nucleocytoplasmic transport is accomplished and regulated by receptor proteins, which bind to the cargo and interact with components of the nuclear pore complexes. In the last few years it was demonstrated that single-molecule microscopy is an excellent and precise tool to study details of cellular nanomachines like the NPC (Peters, 2007). Extensive single-molecule measurements of transport receptors and model substrates have been performed in digitonin-permeabilized cells because this approach avoided the additional complication of in vivo experiments. However, a number of important aspects were certainly absent from in vitro studies compared with the situation in living cells. First, digitonin permeabilization causes a loss of most soluble cytosolic and smaller nuclear components, largely reducing...
overall viscosity and what is known as “molecular crowding” (Ellis, 2001). Because molecular crowding is known to modify binding constants and dynamics (Banks and Fradin, 2005; Hu et al., 2007), it can be anticipated that a complex process such as nucleocytoplasmic transport should behave differently in vivo than in plasma membrane-permeabilized cells. It can also be assumed that such differences should be aggravated by the fact that in permeabilized cells most intranuclear processes are ceased, whereas in living cells processes such as transcription and RNA processing are active; e.g., in experiments analyzing the intranuclear dynamics of the splicing factor U1 snRNP we recently detected clear differences between in vitro and in vivo situations (Kues et al., 2001; Gruenwald et al., 2006b). Sustained transcription and RNA processing will result in continuous nuclear export of mRNA. Export of the usually quite bulky ribonucleoprotein particles could possibly massively impede nuclear import (Naim et al., 2007). Despite the anticipated differences between permeabilized and living cells, the dwell times of all examined molecules were in the same range as those which were recently determined in digitonized cells for the transport receptors p10/NTF2 and kap2 in our laboratory (Kubitscheck et al., 2005), and also for kap1 and a NLS-2xGFP model import substrate by others (Yang et al., 2004; Yang and Musser, 2006). The examined transport receptors, however, belong to quite different transport receptor families. Here we demonstrate a significant robustness of the translocation process with regard to the specific interaction molecular pattern. Musser and co-workers showed that the duration of import substrate interaction with the NPC was a function of the concentration of kap1 in permeabilized cells, and was in the range between 1 and 8 ms for free kap1 concentrations between 15 and 0.1 μM (Yang and Musser, 2006). In living cells we determine dwell times for kap2, kap1, and BSA-NLS of ~7 ms (Table I). Our finding suggests low concentrations of free kap1 in living cells. Only an import-deficient kap1 mutant, kap1ΔN44, and kap2 show a clearly extended interaction time as discussed in the following two paragraphs. All observed dissociation kinetics are predominantly mono-exponential with a minor component showing a greater binding duration in some cases. The fraction of the latter is small (<15%). This is in accordance with previous in vitro measurements. Our results argue for a transport mechanism without major interference between import and export processes, which can be explained by pores being alternatively involved in export or import. It is possible that exporting pores are temporarily unavailable for import, or that indeed two different NPC species exist, one for import or one for export. An alternative scenario would be that NPCs have independent pathways along the NPC channel for import and export (Tran and Wente, 2006). Our measurements, which have been taken under conditions that were as undisturbed as possible, support the notion that molecular crowding caused by ongoing export or further intrinsic cellular components within NPCs in live cells has little effect on translocation times.

The mutant kap1ΔN44 has a RanGTP binding deficiency (Gorlich et al., 1996). In the association time analysis this mutant showed a 1.8-fold extended dwell time at the NPC compared with wild-type kap1 (Fig. 2, Table I). A significantly extended binding time might have been expected, as Ran-GTP binding was suggested to be required for the termination of the import event, i.e., the dissociation of kap2 from kap1 and release of the import substrate from the NPC into the nucleoplasm (Gorlich et al., 1996; Kutay et al., 1997; Sun et al., 2008). However, the ternary complex was not "waiting" for dissociation at the nucleoplasmic exit of the NPC. Rather, our finding is compatible with the assumption that intra-pore movements of transport receptors are persistent random walks along the NPC axis, meaning that kap1 and kap1ΔN44 are free to move within the pore. Kap1ΔN44, however, could not exit the pore at its nucleoplasmic face due to its incapability to bind RanGTP. A Monte Carlo simulation (for details see the online supplemental material, available at http://www.jcb.org/cgi/content/full/jcb.200806173/DC1) of this situation revealed that the residence time within a pore with one “blocked” end, compared with a pore of identical length with both ends open, is indeed prolonged by a factor of 1.75. In this simulation we assumed that the trajectory length inside the pore alone defined the residence time and any further attractive interactions were neglected. Consideration of additional attractive interactions modified the numerical value of prolongation, but the trend of an approximate factor of 2 was maintained. Hence, our result is in good agreement with the notion that RanGTP is required to trigger the release of the ternary import complex on the nucleoplasmic face of the NPC. Certainly, the simple relation between dwell time within the pore and number of "open ends" is only valid under the simplifying assumptions mentioned above. A detailed and sophisticated simulation of binding and transport efficiency of NPCs was recently performed (Zilman et al., 2007).

In some cases (Fig. 2, B, D, and E) mono-exponential dissociation kinetics are not perfectly consistent with the decay data. This observation is in accordance with our previous in vitro results, where we detected small fractions (1–6%) of binding events that were significantly longer than 100 ms, but at that time no attempt was undertaken to analyze these events further. We would like to point out that even a mono-exponential decay does not necessarily imply that the dissociation mechanisms on the nucleoplasmic and cytoplasmic face of the NPC were identical. Generally, only the reciprocal value of the sum of the two—possibly differing—dissociation rates can be determined as a dissociation time constant. Hence, import and export of the transport receptors may well follow different mechanisms with different dwell times, although experimentally only a single dissociation time constant can be monitored. In the present analysis, only kap2 could not be analyzed at all assuming mono-exponential decay (Fig. 2). Here, 10% of the binding events showed a decay time of 100 ms. Previously, we assumed that translocation attempts through clogged NPCs may result in significantly prolonged interactions. Possibly, such events might be the origin of the small, second fraction in the binding duration of kap2. The fact that in the in vivo experiments we observe only moderate increase in the overall number of prolonged binding events compared with experiments in digitonin-permeabilized cells suggests that a strong interference between import and export processes does not occur.

NPCs were marked by POM121-GFP, and the transport receptors and substrates were labeled with the green fluorescent
AlexaFluor 488. The latter were monitored after complete bleaching of the GFP fluorescence in the same fluorescence channel. No additional chromatic corrections or alignments had to be performed, which optimized the colocalization precision. We estimate the error for a single distance measurement to be ± 30 nm (for discussion of errors, see Materials and methods). The distances between the observation sites of the various transport factors to the GFP reference signal were plotted in a histogram representation (Fig. 4). All histograms have an approximate Gaussian shape, and therefore their maximum positions and widths were quantified by fitting Gaussian functions to the data. It is noteworthy that all transport receptors, namely kapβ1, kapα2, and kapβ2, show almost identical widths corresponding to a Gaussian standard deviation of \( \sigma_{exp} = 68 \text{ nm} \) (\( \sigma_{exp} = \text{FWHM}/2.35 \)). A similar distribution was previously obtained for p10/NTF2 (Kubitscheck et al., 2005). These data argue for a related environment for each involved import pathway because NTF2/p10, kapβ1, and kapβ2 belong to independent receptor families and hence pathways. Obviously, the already-discussed robustness of nuclear transport also holds true for the biochemical translocation pathway. As discussed in Materials and methods, the observed width \( \sigma_{exp} \) represented the true spread of binding sites, \( \sigma_a \), convoluted with the colocalization precision, \( \sigma_{coloc} = 30 \text{ nm} \). The true standard deviation of the binding site distribution can therefore roughly be estimated by

\[
\sigma_b = \sqrt{\sigma_{exp}^2 - \sigma_{coloc}^2} = 60 \text{ nm}.
\]

Furthermore, the center of the binding site distribution was in all cases in the range of ±10 nm from the reference site of POM121-GFP. Because the position of POM121 is known to be quite centrally within the NPC (Soderqvist and Hallberg, 1994), transport factors are observed only in the central region of the NPC. Binding occurred neither predominantly in the cytoplasmic nor in the nucleoplasmic filaments because in those cases a wider or even asymmetrical distribution would have been observed. For the same reason, no indication of a putative affinity gradient along the NPC was obtained (Ben-Efraim and Gerace, 2001). The distribution of the import cargo BSA-NLS-Alexa488 was slightly narrower than for the import receptors, corresponding to a binding site distribution width of \( \sigma_{B,\text{BSA}} = 50 \text{ nm} \). Also, the shape of the distribution was more compact, and less blurred compared with those of kapβ1, kapα2, and kapβ2, indicating that the pure geometric distributions of binding sites for ternary complex formation is most realistically described by the histogram of Fig. 4 C.

The binding site distribution of kapβ1ΔN44 was clearly more confined to the center of the pore with \( \sigma_{\text{kapβ1ΔN44}} = 25 \text{ nm} \). The very small value of \( \sigma_{\text{kapβ1ΔN44}} \) indicates a restricted intra-pore motion of kapβ1ΔN44. A free intra-NPC diffusion would extend over a larger distance (Ribbeck and Gorlich, 2002; Yang et al., 2004). Görlich and coworkers detected gold-labeled kapβ1ΔN44 arrested at the NPC, however, ~40 nm from the central plane on the nucleoplasmic side of the NPC (Görlich et al., 1996). In contrast to these reports, Lyman et al. found a complex of GST-IBB-gold and kapβ1ΔN44 accumulating at a distance of 30 nm from the central plane on the cytoplasmic side of the NPC (Lyman et al., 2002). Presumably, details of the methodological approach and data analysis are responsible for these variations. Frey et al. (2006) performed their localization following microinjection into the cytoplasm of live Xenopus oocytes after 2 h incubation, whereas Lyman et al. examined the loci of the transport substrate 30 min after incubation in digitonin-permeabilized HeLa cells. Our in vivo experiments were performed only several minutes after microinjection into the cytoplasm of POM121-GFP expressing HeLa cells. Furthermore, we did not use bulky labels nor bulky substrates.

Numerous problems in biology and biophysics have been studied by single-molecule techniques in the past few years, but the application to living cell systems is still in its infancy (Zlatanova and van Holde, 2006; Ishii and Yanagida, 2007). Single-molecule microscopy and single particle tracking permit a completely new and fascinating insight into biological dynamics (Moerner, 2007). Here we demonstrate that the translocation of proteins across the NPC is autonomous with regard to the respective intracellular physical environment, and robust with regard to the specific biochemical pathway and simultaneously ongoing import–export processes. For the first time, fluorescence single-molecule microscopy provides a direct, real-time insight into molecular processes in living cells with near molecular resolution.

### Materials and methods

#### Cell culture and microinjection

A HeLa cell line stably expressing the GFP-conjugate of POM121 was provided by Brian Burke (University of Florida, Gainesville, FL). HeLa cells were grown in DMEM supplemented with 10% FCS. For live cell analysis, cells were seeded in microscopy dishes (MatTek) up to a confluence of 50%, and examined at room temperature. Microinjection of the probe molecules was performed with an Eppendorf injection and micromanipulation setup using an injection time of 0.8 s at an injection pressure of 100 hPa, and a holding pressure 8 hPa. Before final use of protein probes in microinjection experiments, all probes were diluted and centrifuged at 4 °C, 14,000 rpm for 20 min in transport buffer (50 mM Hepes/KOH, pH 7.3, 110 mM potassium acetate, 5 mM sodium acetate, 2 mM magnesium acetate, 1 mM EDTA, and 2 mM DTT). The concentration of labeled protein in the injection medium was 60 nM for all experiments. Experiments using different types of molecules were performed sequentially to avoid uncertainties about the observed protein species. Different import factors or substrates were never used on the same cell sample.

#### Preparation of import factors and substrate

His-tagged kapα2 and GSTkapβ1 were prepared according to Morociaro et al. (1995). His-tagged karyopherin β2 was purified according to Chook et al. (2002). His-tagged kapβ1ΔN44 was purified as described by Gorlich et al. (1996). BSA was purchased from Sigma-Aldrich, and conjugated to AlexaFluor 488 (Invitrogen) and SV40 large T antigen peptide (Bachem California, Inc.) as described previously (Newmeyer and Wilson, 1991).

#### GST pull-down assay

Functionality of kapα2, kapβ1, and kapβ1ΔN44 after covalent fluorescence labeling with AlexaFluor 488 was confirmed by GST pull-down assays. For this purpose a recombinant import substrate consisting of the SV40-NLS fused to GST was immobilized on Glutathione beads, and then used in a pull-down assay. Therefore, the GST-Sepharose beads were gently washed four times in transport buffer at 4 °C, and 150 μl of beads per sample were centrifuged at 4°C for 3 min at 7,000 rpm. The loaded beads were incubated with either the labeled or unlabeled kapα1 and unbound proteins were removed by gently washing with transport buffer several times. After adding kapα2 and kapα2-488, the beads were incubated for 30 min at 4°C on a rotator and then washed gently four times with transport buffer at 4°C. Beads with bound proteins were dissolved in sample
buffer (200 mM Tris-HCl, pH 6.8, 6% [wt/vol] SDS, 15% glycerin, 10% β-mercaptoethanol, and 0.05% [wt/vol] Bromphenol blue) and resolved on a 10% polyacrylamide gel. As a control, the binding of unlabeled kap2p/kap1p and kap2p/kap1LNA44 to GST-NL5 was analyzed.

**Single-molecule microscopy**

Single-molecule tracking experiments were done with a custom-built single-molecule microscope based on an Axiovert 100TV (Carl Zeiss, Inc.) equipped with a 63X NA 1.4 oil immersion objective lens (for details, see Siebrasse et al., 2007). GFP and AlexaFlour 488 fluorescence was excited by an Ar+ laser emitting at 488 nm. Laser illumination was switched on only during image acquisition by means of an acousto-optical tunable filter. For single-molecule image acquisition an iXon DU 860 BI camera (Andor Technologies) in combination with a 4X magnifier was used. Single-molecule imaging was started 5 min after microinjection of the probes into the cytoplasm to allow cells to recover. Cell recovery after microinjection was monitored by observing the cellular morphology by digitally contrast-enhanced bright field imaging. Then, the green fluorescent signal of the nuclear envelope was used to set the focus at the equatorial plane. Data acquisition was started immediately. After bleaching of the dominant GFP fluorescence, single-molecule signals began to appear in the nucleoplasm, nucleolus and cytoplasm, permitting the direct visualization of the single molecules without tissue sectioning. Generally, 5,000 frames were recorded in a single movie with an integration time of 5 ms, and a frame rate of $k_{ho} = 198$ Hz.

**Image processing of video images**

Images were filtered by the Spot Enhancing filter (Sage et al., 2005) assuming a signal width of 1.8 pixels in Image (Rasband, W.S., Image), National Institute of Health, Bethesda, MD; http://rsb.info.nih.gov/ij/). Filtered and raw image movies were aligned site by site and used for visual identification of signal containing frames. Raw data sequences that contained single-molecule data were separated and later loaded for tracking in Diamtrack 3.0 (Semosaphot, a commercial image processing program for the identification and localization of single particle signals and trajectories (Vallotton et al., 2003). All tracks within a distance of 13 pixels from the image border were discarded. Tracks were stored as text files and analyzed using Origin 7.5 (Microcal).

**Distance determination from NE**

To determine the position of the NE, we fitted the POM-121 signal in Origin 7.5 (Microcal) using a user-defined one-dimensional Gauss fit with a linear ramp (Kubitscheck et al., 2005). The result, a list of NE positions in the NE. Generally, 5,000 frames were recorded in a single movie with an integration time of 5 ms, and a frame rate of $k_{ho} = 198$ Hz.

**Monte Carlo simulation of binding times**

The Monte Carlo model for the binding of labeled proteins (Fig. S1) was tested by immobilization of single proteins on a glass surface, which were imaged under similar conditions as used in the live cell experiments. A Monte Carlo simulation on the effect of release inhibition of the import complex from the NPC was performed to estimate the effect on import complex dwell times at the NPC. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200806173/DC1).

**Online supplemental material**

Photostability of labeled protein (Fig. S1) was tested by immobilization of single proteins on a glass surface, which were imaged under similar conditions as used in the live cell experiments. A Monte Carlo simulation on the effect of release inhibition of the import complex from the NPC was performed to estimate the effect on import complex dwell times at the NPC. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200806173/DC1). The ratio of the binding times was 1:1 in this case. A significantly more detailed and sophisticated simulation of binding and transport efficiency of NPCs was recently performed by Zilm et al. (2007).


